ld 26741-53-7 **Date** 12.12.2003

201-15676B

IUCLID

Data Set

Existing Chemical

: ID: 26741-53-7

CAS No.

: 26741-53-7

EINECS Name

: 2,4,8,10-Tetraoxa-3,9-diphosphaspiro[5.5]undecane, 3,9-bis[2,4-

bis(1,1-dimethylethyl) phenoxy]-

EC No.

: 247-952-5

Molecular Formula

: C33H50O6P2

Status

Memo

: US HPV ULTRANOX 626 Crompton Corp

Printing date

Revision date

: 12.12.2003

Date of last update

: 12.12.2003

Number of pages

: 25

Chapter (profile) Reliability (profile) : Chapter: 1, 2, 3, 4, 5, 6, 7, 8, 10

: Reliability: without reliability, 1, 2, 3, 4

Flags (profile)

: Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

2. Physico-Chemical Data

ld 26741-53-7

Date 12.12.2003

MELTING POINT 2.1

Value

173 - 180 °C

Sublimation

Method

OECD Guide-line 102 "Melting Point/Melting Range"

Year

: 2003

GLP

Test substance

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Product Name: U626 Lot No.: H42265

Result

: No color change was observed during melting point determinations, which

was consistent with U626 being chemically stable at the melting point. : (1) valid without restriction

Reliability 12.12.2003

(14)

2.2 **BOILING POINT**

Value

> 311 °C at 1015 hPa

Decomposition

Method

OECD Guide-line 103 "Boiling Point/boiling Range"

Year GLP

yes

Test substance

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Trade name: U626 Lot No. H42265

Remark

: Estimation using MPBPWIN v 1.40 (US EPA., EPIWIN v 3.10, EPI Suite

Software, 2000) gives a boiling point of 595°C

Result

The duplicate U626 warming curves did not show a distinct boiling plateau up to the maximum temperature of 312°C (at 101.5 kPa). This was supported by the fact that U626 was not observed to boil during the test.

No U626 color change nor smoke was observed during the test.

Reliability

: (1) valid without restriction

12.12.2003

(13)

2.4 **VAPOUR PRESSURE**

Value

.0000000000029 hPa at 25 °C

Decomposition

other (calculated): MPBPWIN v 1.40

Year **GLP**

Method

2003

Test substance

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Test condition

Melting point = 173°C (experimental) Boiling Point = 595 (estimated)

2. Physico-Chemical Data

ld 26741-53-7

Date 12.12.2003

Reliability

(2) valid with restrictions

12.12.2003

(16)

2.5 **PARTITION COEFFICIENT**

Partition coefficient

octanol-water

Log pow

10.9 at °C

pH value

Method

other (calculated): KOWWIN v 1.66

Year

2003

GLP

Test substance

: Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Reliability

: (2) valid with restrictions

12.12.2003

(16)

2.6.1 SOLUBILITY IN DIFFERENT MEDIA

Solubility in

Water

Value

at °C

pH value

concentration

at °C

Temperature effects

Examine different pol.

pKa

at 25 °C

Description

Stable

Deg. product

other: calculated using WSKOW v 1.40

Method Year

2003

GLP

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Remark

An OECD 105 study is currently underway to fill this endpoint.

Result

: Water solubility = 5.67E-8 mg/L

Test condition

Test substance

: Melting point = 173°C (experimental)

Log Kow = 10.9 (estimated)

Reliability 12.12.2003 : (2) valid with restrictions

(16)

3.1.1 PHOTODEGRADATION

Type : air

Light source :

Light spectrum : Nm

Relative intensity : based on intensity of sunlight

INDIRECT PHOTOLYSIS

Sensitizer : OH

Conc. of sensitizer : 1500000 molecule/cm³

Rate constant : .0000000000011 cm³/(molecule*sec)

Degradation : % after

Deg. product

Method : other (calculated): estimation using AOPWIN v1.90

Year : 2003

GLP

Test substance : Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 T1/2 = 1.166 hours

Result : T1/2 = 1.166 hours **Reliability** : (2) valid with restrictions

08.12.2003 (16)

3.1.2 STABILITY IN WATER

 Type
 : Abiotic

 t1/2 pH4
 : at °C

 t1/2 pH7
 : at °C

 t1/2 pH9
 : at °C

 Deg. product
 : No

 Method
 : OECD 111

Year : OECD 11°

GLP :

Test substance : 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-bis[2,4-bis(1,1-

dimethylethyl)phenoxy]-CAS No.: 26741-53-7

Purity: >96%

Source: General Eletrical

Method : The test substance was added to pH 4, 7 and 9 buffered water and

incubated at 25°C. Samples were analyzed after Day 1, 2, 3, 4, 6, 8 and

10 and analyzed by HPLC.

Result : The test substance was practically insoluble in water and difficult to

determine the hydrolytic half-life. However, it is expected to be

hydrolytically unstable similar to other organophosphite class of chemicals

at all pHs'.

Fugacity Model Half-Life: 3.6 e003 hr

Reliability : (1) valid without restrictions

10.21.2004 (14)

ld 26741-53-7 **Date** 12.12.2003

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type : fugacity model level III

Media

Air : % (Fugacity Model Level I)

Water : % (Fugacity Model Level I)

Soil : % (Fugacity Model Level I)

Biota : % (Fugacity Model Level II/III)

Soil : % (Fugacity Model Level II/III)

Method : other: EPIWIN Level III Fugacity Model

Year : 2003

Test Substance : Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Test condition : Henry's Law Constant: 3.24E-9 atm-m3/mol (Henrywin)

Vapor pressure: 2.9E-12 mmHg (Mpbpwin)

Melting Point: 173°C (experimental)

Log Kow: 10.9 (Kowwin)

Soil Koc: 3.26E10 (calc by model)

	Mass Amount	Half-life	Emissions	
	(percent)	(hr)	(kg/hr)	
Air	0.0175	2.33	1000	
Water	1.25	3600	1000	
Soil	33.1	3600	1000	
Sediment	65.6	14400	0	

	Fugacity (atm)	Reaction (kg/hr)	Advection (kg/hr)	Reaction (percent)	Advection (percent)
Air	1.12E-16	881	29.6	29.4	0.988
Water	1.07E-19	40.8	212	1.36	7.06
Soil	2.14E-21	1080	0	36	0
Sediment	1.9E-19	535	222	17.8	7.41

Persistence time: 5640 hr Reaction time: 6680 hr Advection time: 36500 hr Percent reacted: 84.5 Percent advected: 15.5

Half-lives (hr), (based upon Biowin (ultimate) and Aopwin):

Air: 2.33 Water: 3600 Soil: 3600 Sediment: 14400 Biowin estimate: 0.802

Advection times (hr):

Air: 100 Water: 1000 Sediment: 5E+4

Reliability : (2) valid with restrictions

08.12.2003

(16)

3.5 BIODEGRADATION

Type

aerobic

Inoculum

Deg. product

Method other: Estimation using BIOWIN v4.00 Year 2003

GLP

Test substance Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Result : MITI Linear Biodegradation Probability: -0.6156

MITI Non-linear Biodegradation Probability: 0.000

The substance is predicted to be not readily biodegradable

Reliability

: (2) valid with restrictions

12.12.2003

(16)

4.1 **ACUTE/PROLONGED TOXICITY TO FISH**

Type other: estimation

Species

Exposure period 96 hour(s)

Unit : mg/l LC50 1.93E-6

Method other: calculated using ECOSAR v 0.99g

Year 2003

GLP

Test substance : Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Reliability : (2) valid with restrictions

12.12.2003 (16)

4.2 **ACUTE TOXICITY TO AQUATIC INVERTEBRATES**

Type other: estimation

Species Daphnia sp. (Crustacea)

Exposure period : 48 hour(s) Unit mg/l EC50 3.82E-6

Method other: calculated using ECOSAR v 0.99g

Year

GLP

Test substance : Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Reliability : (2) valid with restrictions

12.12.2003 (16)

4.3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE

Species : Endpoint :

 Exposure period
 : 96 hour(s)

 Unit
 : mg/l

 EC50
 : 3.99E-6

Method : other: calculated using ECOSAR v 0.99g

Year : 2003

GLP :

Test substance: Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Reliability : (2) valid with restrictions

12.12.2003 (16)

5.1.1 ACUTE ORAL TOXICITY

Type : LD50

Value : 5580 mg/kg bw

Species : rat

Strain : Sprague-Dawley
Sex : male/female

Number of animals : 100

Vehicle

Doses : 1300 - 15300 mg/kg Method : other:IBTL method

Year : 1975 **GLP** : No

Test substance: Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Product Name: Ultranox 626 (Weston 6140)

Reliability : (4) not assignable

12.12.2003 (4)

Type : LD50

Value : > 6080 mg/kg bw

Species : hen
Strain : Leghorn
Sex : female
Number of animals : 24

Vehicle : other: corn oil

Doses : 800, 1200, 1800, 2700, 4050, 6080 mg/kg bw

Method : other: Biodynamics Inc method

 Year
 : 1980

 GLP
 : No

ld 26741-53-7

Date 12.12.2003

Test substance

Chemical name: 2.4.8.10-tetraoxa-3.9-diphosphaspiro [5.5]undecane, 3.9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Trade name: Weston 1452

Purity: 99.5% Lot No.: PP-357

Method

No of animals/sex/dose: 4 hens/dose

Vehicle: Corn oil

Route of administration: Gavage

Result

: Number of deaths at each dose level: No hens died during the study

Clinical signs: Occasional hens showed green or yellow discoloration of the feces on the day of dosing. This reflects the use of corn oil vehicle and is not considered to indicate a response to treatment. There were no signs of toxicity and the hens remained normal in respect of appearance, mood. locomotor function and body weight throughout the observation period.

Necropsy findings: Occasional findings of liver discoloration. This is a common pathological entity in hens of this age and is not considered to

reflect any delayed or residual response to treatment.

Test condition

: Age: 12 months

Weight: 1.3 - 1.9 kg

Volume administered: 10 mL/kg Post dose observation period: 14 days

Reliability

: (2) valid with restrictions 08.12.2003

(1)

5.1.2 ACUTE INHALATION TOXICITY

Type

LC50

Value

 $> 2000 \text{ mg/m}^3$

Species

Strain

Sprague-Dawley

Sex

male/female

Number of animals

10

Vehicle

Doses

2 ma/L : 1 hour(s)

Exposure time **Test Substance**

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Trade name: weston 61440

Result

None of the animals died and all the animals appeared normal during the 14 day oservation period. The gross pathological autopsy of all 10 animals

showed normal appearance of the organs of the thorax and the abdomen. : (4) not assignable

Reliability

12.12.2003

(11)

5.1.3 ACUTE DERMAL TOXICITY

Type

LD50

Value

: > 2000 mg/kg bw

Species : rabbit

Strain : New Zealand white

Sex : male/female

Number of animals : 10

Vehicle

Doses : 2000 mg/kg bw

Method : OECD Guide-line 402 "Acute Dermal Toxicity"

Year : 1994 **GLP** : yes

Test substance: Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Trade name: Ultranox 626

Lot No.: HCC033

Result: Mortality: There were no deaths during the study

Clinical observations: 5 rabbits had abnormal defecation (soft stool, mucoid feces) on day 0 or 1. One of these animals also had wet, and subsequently, dry brown urogenital matting. These findings were considered to be a result of the bandage/restraint procedures used and not related to the test material. An additional spontaneaous occurrence of soft stool was noted on day 9 for one animal. There were no other findings.

Dermal observations: The test material induced generally very slight to slight erythema on all rabbits and very slight to slight edema on 8 rabbits. There was a single occurrence of moderate erythema at the day 1 observation. Desquamation was present on five sites by day 7. There were no other dermal findings. One site had very slight erythema at study termination (day 14).

Body weights: There wer no remarkable changes or differences noted in body weights during the study.

Necropsy: Accessory splenic tissue, a common congenital abnormality in this strain of rabbit, was noted for four animals at the terminal necropsy. There were no other gross necropsy findings for all examined tissues.

Test condition : Age: Approximately 11 weeks old

Weight: 2014 - 2224 g at study initiation Post dose observation period: 14 days

Reliability : (1) valid without restriction

Guideline study conducted to GLP

28.11.2003 (10)

5.2.1 SKIN IRRITATION

Species: rabbitConcentration: .5 gExposure: OcclusiveExposure time: 24 hour(s)

Number of animals : 6

Vehicle : physiol. saline

PDII

Result : corrosive

ld 26741-53-7 Date 12.12.2003

Classification

Method

: other: 16CFR 1500.42

Year

1981

GLP

no

Test substance

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Trade name: Weston XP-1532

Lot No.: 1198

Result

: Necrotic skin occurred in all animals at 24 and 72 hours. The responses

for abraded and intact skin were the same.

Reliability

12.12.2003

: (2) valid with restrictions

(3)

5.3 **SENSITIZATION**

Type

other: Footpad method

Species

guinea pig

Concentration

Induction 1 % other: injected into footpad 2nd: Challenge 10 % open epicutaneous

3rd:

Number of animals

: 20

Vehicle

other:Induction: Freund's adjuvant, Challenge: acetone/dioxane/guinea pig

fat 7:2:1

Result

not sensitizing not sensitizina

Classification Method

OECD Guide-line 406 "Skin Sensitization"

Year

GLP

: ves

Test substance

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Trade name: Ultranox 626

Lot No.: U626-1

Result

Primary irritation screen: No signs of erythema or edema were evident for

the animals that were administered 1, 3 or 10% of the test material.

Sensitization study: No signs of erythema or edema were noted at the 24 or 48 hour for animals previously induced with either Freund's adjuvant (control animals) or with 1% of the test material in Freund's adjuvant).

No toxic effects or systemic clinical signs were ntoed during the study.

All animals gained weight normally during the study.

Test condition

: Strain: Crl:(HA)BR VAF/PlusT

Primary irritation screen:

No. of animals: 5/dose Doses: 1.0%, 3.0%, 10.0% Sex: Not determined

Body weight range: 571 - 741 g

Age: Not determined

ld 26741-53-7 **Date** 12.12.2003

Sensitization study:

No. of animals: 20; 10 control, 10 test

Sex: Female

Body weight range: 361 - 442 g Age at study initiation: 6 - 7 weeks

Reliability

: (1) valid without restriction

Guideline study conducted to GLP

28.11.2003

(15)

5.4 REPEATED DOSE TOXICITY

Type

.

Species

Rat

Sex

male/female

Strain

: other: Charles River CD

Route of admin. Exposure period

: oral feed

Frequency of treatm.

: 90 days : Daily

Post exposure period

None

Doses

0, 100 ppm; 300 ppm; 1000 ppm

Control group

: yes, concurrent vehicle

NOAEL Method 300 ppm (Males: 22 mg/kg/bw; Females 26 mg/kg bw)

: other: IRDC method

The test substance was fed in the diet to three groups of 20 male and 20 female rats. The rats were observed daily for signs of overt toxicity and mortality. Detailed observations including incidence, size and location of palpable masses and individual body weights were recorded weekly. Individual food consumptions were recorded daily. All rats received the ophthalmic examinations during the pretest period and at 3 months of study. Clinical laboratory test were performed for 10 rats/sex/group at 1 and 3 months of study. At the termination of the study period, selected organ weights from 10 male/10 female rats were recorded and tissues (i.e., adrenals, aorta, eye, stomatch, liver, spllen, pancreas, urinary baldder, sternum, prostate/uterus, testes/ovaries, brain, heart, lung, pituitary, mammary gland, thymus, kidneys, etc) were examined.

Year

1979

GLP

No

Test substance

: Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Trade name: Weston XR-1532

Lot No.: 225-35

Impurities: 1% tris-isopropanolamine

Result

Body weight: The group mean body weights of both the low and mid dose male and female rats were greater than control values, and both groups of high dose males and females had group mean body weights lower than

controls.

Food/water consumption: Low and mid dose male rats consumed slightly more food than did controls, while high dose males ate slightly less than control. All of the groups of females consumed approximately equivalent

amounts of food.

Clinical signs: No signs of overt toxicity were observed among the treated rats. Incidental signs observed for a few control and/or treated rats included hair loss. missing or malaligned upper incisor, excessive lacrimation, red material around the eyes, leaning to left, red or swollen eyes, rales, corneal opacity, swollen ventral neck, swollen conjunctiva and dilated pupil.

Ophthalmologic findings: There were no compound related effects noted during the 3-month ophthalmic examinations.

Hematology: No compound related effects on the results of the hematologic tests were observed.

Clinical biochemistry: No compound related effects on the results of the biochemical tests were observed.

Mortality: No compound related effects on survival rates were noted (one female rat in the 1000 ppm group died during the study).

Gross pathology: No compound related gross lesions were observed in any of the rats from the treated groups. No tumors were noted upon gross examination.

Organ weight changes: Statistically significant variations (p<0.01) in absolute weights of kidneys of male rats at 300 ppm and hearts of male rats at 100 and 300 ppm were not considered compound related.

Histopathology: Microscopic lesions considered probably compound

related were seen in livers and spleens of the female rats from the 1000 ppm group. This consisted of very slight to slight extramedullary hematopoiesis in these organs. This lesion was not present in rats from the control and the 300 ppm groups but was seen in one rat from the 100 ppm group. Other microscopic lesions in livers and those seen in other organs in the control and 1000 ppm groups were considered spontaneous in nature and unrelated to the administration of the compound. Statistical methods: All statistical analyses compared the treatment groups with the control group by sex. Body weights, food consumption, absolute and relative organ weights and hematology, biochemistry and urinalysis parameters were compared by analysis of variance (one-way), Bartlett's test for homogeneity of variances and the appropriate t-test (for equal or unequal variances) as described by Steel and Torrie using Dunnett's multiple comparison tables to judge significance of differences.

Test condition

Test subjects:

Weight at study initiation: 80-107g (male), 77-98g (female)

No. of animals/sex/dose: 20 male, 20 female

Study Design:

Vehicle: Feed (Rodent Laboratory Chow)

Clinical observations performed and frequency: The rats were observed twice daily for signs of overt toxicity and mortality. Detailed observations were recorded weekly and included the size, incidence and location of all

ld 26741-53-7

Date 12.12.2003

palpable masses. Individual body weights were recorded weekly and individual food consumption was recorded daily.

Organs examined at necropsy: The following tissues from 10 males and 10 females from the control group and the 1000 ppm group were examined microscopically: adrenals, aorta, eye and optic nerve, esophagus, stomach, duodenum, jejunum, ileum, cecum, colon, rectum, liver, trachea, spleen, pancreas, urinary bladder, bone marrow (sternum), prostate/uterus, seminal vesicles, testes/ovaries, brain, heart, lung and bronchi, sciatic nerve, pituitary, thyroid and parathyroid, mesenteric lymph node, mandibular lymph node, spinal cord, salivary gland (submaxillary), skeletal muscle (thigh), skin, mammary gland, thymus, kidneys, any other tissue with gross lesions.

Bone marrow smears from all rats were made at necropsy and examined microscopically.

Additionally, livers and spleens from 10 male and 10 female rats from the 100 ppm and 300 ppm groups were examined.

Reliability

: (2) valid with restrictions

Well conducted study, prior to GLP

05.12.2003

(9)

Туре

Dog

Species Sex Strain

: male/female : Beagle

Route of admin.

other: oral via gelatin capsule

Exposure period
Frequency of treatm.
Post exposure period

4 months daily

Doses

: 0, 4, 12, 40 mg/kg/day

Control group

yes

NOAEL

12 mg/kg bw

Method

other: Bio/dynamics Inc method

Year GLP 1980

Test substance

Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5,5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Trade name: Weston XR-1532 Lot No.: 1532-003-04189-X

Purity: 100%

Result

Mortality: One high dose female was sacrificed in a moribund condition after 86 days on test. All of the remaining control and treated animals survived the duration of the study.

Physical and neurological observations: The only physical and neurological observations noted during the study which were considered related to the administration of the test substance involved the high dose female which was sacrificed in a moribund condition. On test day 51, this animal exhibited the first signs of uncoordination of the hind limbs and a poor righting reflex. These symptoms progressed and by test day 58, the animal displayed a lack of coordination of both the fore and hind limbs. The

animal became progressively recumbent as the uncoordination progressed to rigid fore and hind limb paralysis. During this time the animal was offered canned dog food in an attempt to increase its food consumption and maintain its body weight. However this was unsuccessful and the animal was sacrificed in a moribund condition on test day 86.

Ophthalmology: There were no ocular abnormalities observed after 3 or 4 months of study which could be attributed to the test substance.

Body weight and food consumption: The mean body weights of all treated males were slightly greater than control prior to initiation of dosing (4-5%) and throughout the 4 month treatment period. At 18 weeks these differences from control were 25%, 20%, and 21% in the low, mid and high dose males, respectively. Mean food consumption values were slightly lower in the low (13%), mid (12%) and high (17%) dose males when compared to the control values over the 18 week period of test substance administration. Mean body weight and food consumption values for the control and treated females were unremarkable throughout the study with the exception of one high dose female. During week 9 this animal began to exhibit decreases in food consumption and body weight. These decreases continued over the next 3 weeks and the animal was sacrificed in a moribund condition on test day 86.

Hematology and clinical chemistry: At 1 month the mean platelet counts of the treated males were significantly lower than control. However, these differences were attributed to a slightly greater mean platelet counts in the control group rather than an effect of the test substance. Other differences from control were noted in some of the hematology and clinical chemistry parameters evaluated. However, these differences were not dose related or consistent over time and therefore were not considered related to administration of the test material.

Urinalysis: The urinalysis data collected after 1, 3 and 4 months of study were unremarkable and revealed no differences between control and treated groups which could be attributed to the administration of the test substance.

Organ weights and organ/body weight ratios: Slight diferences from control, some statistically significant, were observed in the mean absolute and relative organ weights of the treated animals. These variations were attributed to differences in terminal body weights and normal biological variation between animals. There were no differences observed in absolute or relative organ weights which were attributed to the administration of the test substance.

Pathology: A subacute, eosinophilic pneumonia was observed microscopically in four of eight high dose animals, one of eight mid dose animals and two of eight low dose animals. This condition was not observed in any of the control animals. The etiology of the pulmonary lesions could not be determined. Seven of the eight high dose animals displayed degenerative myelin lesions. These lesions were confined to the high dose group and were considered related to the administration of the test material. One animal (the female which died) displayed clinical manifestations of several abnormalities of the axonal fibers and myelin. Statistical methods: Hematology and clinical chemistry: Test substance groups were compared to control by the F-test and Student's T-test. When

variances differed significantly (F-test), Student's t-test was appropratiately modified using Cochrane's approximation (t'). Snedecor, G.W. & Cochran, W.G. (1967) Statistical Methods, 6th ed. Iowa State University Press, Ames, pp 104-106, 114-119. Body weight, Food consumption, Organ weights and organ/body weight ratios: Treated groups were ocmpared to control. Dunnett, C.W. (1964) J. Am. Stat. Assn. 50, 1096-1121 and Biometrics 20, 482 (1964).

Test subjects:

Weight at study initiation: 80-107g (male), 77-98g (female)

No. of animals/sex/dose: 4/sex/dose

Study Design:

Vehicle: None

Clinical observations performed and frequency: Twice daily for mortality and gross signs of toxicological or pharmacologic effects.

Detailed physical examination: Pretest and weekly thereafter

Neurologic examination: Pretest and monthly thereafter

Ophthalmoscopic examination: Pretest, 3 and 4 months

Body weight: Pretest, weekly during treatment and terminally

Food consumption: Pretest and weekly thereafter

Laboratory studies: Pretest and monthly thereafter. Parameters evaluated - Hematology: hemoglobin, hematocrit, erythrocytes, platelets, clotting time, prothrombin time, total and differential leukocytes, erythrocyte morphology. Clinical chemistry: serum glutamic oxaloacetic transaminase, serum glutamic pyruvic transaminase, alkaline phosphatase, bliid urea nitrogen, fasting glucose, total protein, albumin, globulin, A/G ratio, cholesterol, triglycerides, sodium, potassium, chloride, calcium, creatinine, lactic acid dehydrogenase, total bilirubin, direct bilirubin. Urinalysis: gross appearance, specific gravity, pH, protein, glucose, ketones, bilirubin, occult blood, microscopic analysis

Organs examined at necropsy: Organs weighed and organ/body weight ratios calculated: adrenals, brain, ovaries, testes (without epididymides), heart, kidneys, liver (with gall bladder attached), pituitary, spleen, thyroid (with parathyroid). Tissues examined histopathologically: adrenal, aorta (thoracic, arch, lumbar), bone and bone marrow (sternum), brain, epididymus, esophagus, eye, heart, intestine, cecum, colon, rectum, kidney, liver with gall bladder, lungs with mainstem bronchi, lymph nodes (cervical, peribronchial, mesenteric), nerve (left sciatic), oral mucous membrane (including tongue, buccal, maxillary gingiva, pharynx and nasopharynx), ovary, pancreas, pituitary, prostate, salivary gland (submaxillary), skeletal muscle (biceps femoris), skin with mammary gland (left inguinal), spinal cord (cervical, thoracic, lumbar), spleen, stomach (cardia, pylorus, fundus), testis, thymus, thyroid with parathyroid, trachea,

urinary bladder, uterus (corpus, cervix, fallopian tubes), vagina, gross lesions, tissue masses or suspect tumors and regional lymph nodes.

(2)

Reliability : (2) valid with restrictions

Well conducted study, prior to GLP

05.12.2003

Туре

Species : Rat

Sex : Male/female
Strain : Wistar
Route of admin. : oral feed
Exposure period : 24 months
Frequency of treatm. : Daily
Post exposure period : None

Doses : 100, 500 ppm

Control group : yes, concurrent no treatment

NOAEL : 500 ppm

Method : other: NIHMR method

Year : 1983 GLP : No

Test substance : Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Trade name: Weston XP1452

Purity: 99%

Impurities: Tri-isopropanolamine (1%)

Result : Body weight: No diference was observed between the average weights of

the control group and treated animals.

Food/water consumption: No difference was observed between the treated groups and the control group animals.

Clinical signs: No difference between the treated animals and the control group. Toward the 18th month, the animals began to show signs of ageing, i.e. some of them had a yellow coat, dyspnoea, pulmonary insufficiency and dark rings around the eyes, but these symptoms were relatively rare and no more frequent among the treated animals than among the control animals.

Hematologic findings: The haemoglobin level, the haematocrit, the number of erythrocytes and the level or relative number of leucocytes were the same in all the groups.

Clinical biochemistry: No difference in glucose levels, ureic nitrogen levels, the GOT, the GPT, the PA, total proteins and the albumin levels were seen.

Renal function: The urine density showed no significant variation and neither did the proteinurea, glycosuria and urinary residue examinations.

Mortality:

Dose Sex Total Months to death (mg/kg) Deaths (No. of animals)

ld 26741-53-7

Date 12.12.2003

0	Male	11	13(1), 19(4), 20(1), 21(3), 22(2)
	Female	5	20(1), 21(2), 22(2)
100	Male	9	12(1), 13(1), 15(1), 16(1), 17(1), 20(1). 21(2), 22(1)
	Female	5	19(1), 21(3), 22(1)
500	Male	15	13(1), 14(2), 15(1), 16(1), 17(3), 18(1), 20(1), 21(1), 22(4)
	Female	6	14(2), 15(1), 21(1), 23(1)

Mortality was higher among the males than among the females. A dose of 100 ppm did not give rise to higher mortality amog the male or female rats compared to the control group. At 500 ppm, mortality is no higher among the female rats, but higher mortality was observed among the male rats. The authors concluded that this difference was not significant.

Gross pathology: an examination of the main organs did ot reveal any significant difference between the treated rats and the control group. The animals often showed clear organic pathological signs of aged animals which could in no case be attributed to the treatment since they appeared with equal frequency in the control group.

Organ weight changes: No change in the weight of the main organs under the effect of the treatment was seen.

Histopathology: No greater frequency of anomalies in the organs of the treated rats compared with the control group.

Test subjects

Age at study initiation: no data

No. of animals/sex/dose: 30 males/females per dose

Study Design

Vehicle: Feed

Clinical observations: Each animal was weighed every week over the first four weeks, twice a week during weeks 6-12 and then every other week after week 12. Food and water intake was measured in weeks 1+3, 3+4, 11+12, 23+24, 38+39, 52+53, 75+76, 86+87 and 97+98.

Haematological examinations were conducted during weeks 12, 24, 52, 88 and 104. The blood was used:

- a) to conduct measurement of haemoglobin content, haematocrits, number of erythrocytes and leucocytes and leucocytic formula;
- b) to determine blood glucose and ureic nitrogen levels;
- c) to analyze for the following enzymatic activities: glutamic-pyruvate transaminase, alkaline phosphatase and proteins and albumin serum.

Urine was collected in weeks 12, 24, 53, 98-104 and tested for appearance, pH, glucose, proteins, density, blood, lymphocytes and erythrocytes, epithelial cells, urate and phosphate crystals and evaluation of the bacterial population.

Test condition

ld 26741-53-7 Date 12.12.2003

Organs examined at necropsy: The main oragans were subjected to a macroscopic examination and some were weighed (liver, kidneys, splen. brain, heart, supra-renal bodies and hypophysis). Tissue fragments were taken: heart, kidney, liver, pancreas, spleen, brain, ovary or testicle, thyroid, suprrenal bodies, thymus, lung, trachea, salivary glands, stomach, duodenum, colon, caecum and uterus.

: (2) valid with restrictions

Reliability 10.12.2003

(5)

i.p GENETIC TOXICITY 'IN VITRO'

Type

: Ames test

System of testing

Salmonella typhimurium strains TA97, TA 98, TA100, TA102

Escherichia coli Strain WP2 (PKM101)

Test concentration

50 - 2000 μg/Plate

Cycotoxic concentr. Metabolic activation

: with and without

Result

: negative

Method

: other: Hachiya (1994)

Year **GLP**

: 1994

Test substance

: no data : Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7

Test condition Reliability

: Solvent: Acetone and Tween 80

(4) not assignable

Secondary literature data

08.12.2003

(8)

Type

Chromosomal aberration test

System of testing

Chinese hamster ovary cells

Test concentration

31.3-2000 µg/mL (-S9, 4 hour exposure), 31.3 - 3000 µg/mL (+S9, 4 hour

exposure; -S9, 20 hour exposure) : 500 µg/mL (+S9), 2000 µg/mL (-S9)

Cycotoxic concentr. Metabolic activation

with and without

Result

positive

Method

OECD Guide-line 473

Year **GLP**

: yes

Test substance

2003

: Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3.9bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

> CAS No.: 26741-53-7 Trade name: Ultranox 626

Lot No.: H42265

Method

: METHOD

Metabolic activation: Arochlor 1254-induced rat liver S(9)

Concentrations tested:

Experiment 1:

31.3, 62.5, 125, 250, 500, 1000 µg/mL (-S9, 4 hour exposure) 31.3, 62.5, 125, 250, 500, 1000, 1500, 3000 µg/mL (-S9, 20 hour

ld 26741-53-7

Date 12.12.2003

exposure)

31.3, 62.5, 125, 250, 500, 1000, 1500, 3000 (+S9, 4 hour exposure)

Statistical Methods: Statistical analysis of the percent aberrant cells was performed using Fisher's exact test. Fisher's exact test was used to compare pairwise the percent aberrant cells of each treatment group with that of the solvent control. In the event of a positive Fisher's exact test at any test article dose level, the Cochran-Armitage test was used to measure dose-responsiveness.

Result

- +S9, 4h exposure: The percentage of cells with structural aberrations in the test article treated group was statistically increased above that of the solvent control at 250 μ g/mL (p=0.05, Fisher's exact test). The Cochran-Armitage test was also positive for a dose response (p<0.05). However, the percentage of cells with structural aberrations in the test article treatd group (6.0%) was within the historical solvent control range of 0.0 6.5%. Therefore it is not considered to be biologically significant. The percentage of cells with numerical aberrations in the test article treated groups was not significantly increased above that of the solvent control at any dose level (p>0.05, Fisher's exact test).
- -S9, 20h exposure: The percentage of cells with structural or numerical aberrations in the test article treated groups was not significantly increased above that of the solvent control at any dose level (p>0.05, Fisher's exact test).
- -S9, 4h exposure: Due to the lack of dose levels with =5-% toxicity relative to solvent control in this group, the experiment was repeated at higher dose levels
- –S9, 4h exposure (repeat): The percentage of cells with structural aberrations in the test article treated groups was significantly increased above that of the solvent control at 2000 μg/mL (p=0.05, Fisher's exact test). The Cochran-Armitage test was also positive for a dose related response (p<0.05). The percentage of cells with numerical aberrations in the test article-treated groups was not significantly increased above that of the solvent control at any dose level (p=0.05, Fisher's exact test). Test Design

Test condition

Number of replicates: Duplicate tests

Positive and negative controls and treatment: Mitomycin C was used as the positive control in the non-activated study at final concentrations of 0.1 and 0.2 μ g/mL. Cyclophosphamide was used as the positive control in the S9 activated study at final concentrations of 10 and 20 μ g/mL. The solvent for the test article (DMSO) was used as the solvent control at the same concentration as that found in the test article-treated groups.

Solvent: dimethylsulfoxide

Number of metaphases analyzed: Whenever possble, a minimum of 200 metaphase spreads (100 per duplicate flask) were examined and scored for chromatid-type and chromosome-type aberrations. The number of metaphase spreads that were examined and scored per duplicate flask was reduced when the percentage of aberrant cells reached a statistically significant level before 100 breaks were scored.

Description of follow up repeat study: Due to lack of dose levels with 50% toxicity relative to the solvent control in the non-activated 4 hour exposure group, the chromosome aberration assay was repeated in that group at dose levels of 250, 500, 1000, 1250, 1500 and 2000 µg/mL.

Criteria for evaluating results: The test article was considered to induce a positive response when the percentage of cells with aberrations increased in a dose-responsive manner with one or more concentrations being statistically significant (p=0..5). However, values that were statistically significant but did not exceed the range of historic solvent controls was judged as not biologically significant. Test articles not demonstrating a statistically significant increase in aberrations were concluded to be negative.

Conclusion

: Under the conditions of the assay described in this study, the test material was concluded to be weakly positive for the induction of structural and negative for the induction of numerical chromosome aberrations in CHO cells in the absence of metabolic activation. The test material was concluded to be negative for the induction of structural and numerical chromosome aberrations in CHO cells in the presence of metabolic activation.

(7)

Reliability 25.11.2003

: (1) valid without restriction

(1) Valid Williagt Toolifolion

i.p GENETIC TOXICITY 'IN VIVO'

Type : Micronucleus assay

Species : mouse Sex : male/female

Strain : ICR Route of admin. : i.p.

Route of admin. : i.p. Exposure period : 24, 48 hours

Doses : 500, 1000, 2000 mg/kg

Result : negative

Method : OECD Guide-line 474 "Genetic Toxicology: Micronucleus Test"

Year : 2003 **GLP** : yes

Test substance : Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Trade name: Ultranox 626

Lot No.: H42265

Result : Effect on mitotic index or PCE/NCE ratio by dose level by sex: See table

below

Genotoxic effects: Negative

Mortality at each dose level by sex:

Pilot toxicity study: No mortality occurred at any doses during the course of

the study.

Main study: No mortality occurred at any dose during the course of the

micronucleus study.

Clinical signs:

Pilot toxicity study: All animals appeared normal except male mice at 2000 mg/kg, which exhibited piloerection.

Main study: Piloerection was seen in all male and female mice at 1000 and 2000 mg/kg. All other mice appeared normal during the study.

Bodyweight changes:

Pilot toxicity study: Change in group mean bodyweights ranged from -0.3% (1.0 mg/kg) to 4.5% (1000 mg/kg) after 3 days.

Mutant/aberration/mPCE/polyploidy frequency, as appropriate: See table below.

Food/water consumption: no data

Age at study initiation: 6 - 8 weeks old at the initiation of each phase of the study.

No. of animals per dose:

Pilot toxicity study: 2 male mice dosed at 1, 10, 100 or 1000 mg/kg b.w.; 5 male and 5 female mice dosed at 2000 mg/kg.

Main study: Groups of 5 male/5 female mice dosed at 0, 500, 1000, 2000 mg/kg (euthanized at 24 h); Groups of 5 male/5 female dosed at 0, 2000 mg/kg (euthanized at 48 h).

Route: i.p.

Vehicle: Corn oil.

Controls: Vehicle (Corn oil), cyclophosphamide monohydrate (positive).

Clinical observations performed: Clinical signs, mortality, bodyweight

Organs examined at necropsy: none

Criteria for evaluating results: The incidence of micronucleated polychromatic erythrocytes per 2000 polychromatic erythrocytes was determined for each mouse and treatment group. Statistical significance was determined using the Kastenbaum-Bowman tables which are based on the binomial distribution. In order to quantify the proliferation state of the bone marrow as an indicator of bone marrow toxicity, the proportion of polychromatic erythrocytes to total erythrocytes was determined for each animal and treatment group. The test article was considered to induce a positive response if a dose-responsive increase in micronucleated polychromatic erythrocytes was observed and one or more doses were statistically elevated relative to the vehicle control (p<=0.05, Kastenbaum-Bowman Tables) at any sampling time. However, values that were statistically significant but did not exceed the range of historical negative or vehicle controls were judged as not biologically significant. The test article was judged negative if no statistically significant increase in micronucleated polychromatic erythrocytes above the concurrent vehicle control values and

Test condition

ld 26741-53-7

Date 12.12.2003

no evidence of dose responses were observed at any sampling time.

Criteria for selection of M.T.D.: based on preliminary toxicity study.

Table: Summary of Bone Marrow Micronucleus analysis

Treatment	Sex	Time	No. of	PCE/Total	Change from	Micronucleated Polychromatic Erythrocytes	
(20mL/kg)		(hr)	mice	Erythrocytes	Control (%)	Number per 1000 PCEs Number p	
				(mean ± SD)		(mean ± SD)	PCEs Scored ¹
Corn oil	M	24	5	0.456±0.07	-	0.6±0.22	6/ 10000
	F	24	5	0.526±0.09	-	0.5±0.35	5/ 10000
Test article							
500 mg/kg	M	24	5	0.477±0.06	5	0.6±0.22	6/ 10000
	F	24	5	0.462±0.04	-12	0.6±0.42	6/ 10000
1000 mg/kg	M	24	5	0.480±0.06	5	0.7±0.27	7/ 10000
	F	24	5	0.543±0.05	3	0.8±0.27	8/ 10000
2000 mg/kg	M	24	5	0.493±0.09	8	0.5±0.00	5/ 10000
	F	24	5	0.469±0.04	-11	0.3±0.27	3/ 10000
CP ²	M	24	5	0.335±0.03	-27	22.2±2.20	*222/ 10000
50 mg/kg	F	24	5	0.325±0.01	-38	20.4±2.43	*204/ 10000
Corn oil	M	48	5	0.502±0.06	-	0.3±0.27	3/ 10000
	F	48	5	0.483±0.05] -	0.6±0.22	6/ 10000
Test article	Test article						
2000 mg/kg	M	48	5	0.447±0.05	-11	0.6±0.22	6/ 10000
	F	48	5	0.495±0.05	2	0.5±0.35	5/ 10000

^{1*}statistically significant, p<=0.05 (Kastenbaum-Bowman Tables).

Reliability

: (1) valid without restriction

26.11.2003

(6)

5.8.1 TOXICITY TO FERTILITY

5.8.2 DEVELOPMENTAL TOXICITY/TERATOGENICITY

Species Sex

: rabbit

: female

Strain

: oral unspecified

Route of admin. Exposure period

Days 6 to 18 of gestation

Frequency of treatm.

: Daily

Duration of test

: Fetuses removed on day 29 of gestation

Doses

: 0, 20, 50, 200 mg/kg

Control group

: Yes

Method

The test material was administered to 4 groups of 15 rabbits at dose levels

of either 0, 20, 50 or 200 mg/kg from days 6 to 18 of gestation. The fetuses were removed by caesarean section on day 29 of gestation and

examined for malformations.

Year **GLP**

Well conducted study, prior to GLP

² cyclophosphamide monohydrate

ld 26741-53-7

Date 12.12.2003

Test substance

: Chemical name: 2,4,8,10-tetraoxa-3,9-diphosphaspiro [5.5]undecane, 3,9-

bis[2,4-bis(1,1-dimethylethyl)phenoxy]-

CAS No.: 26741-53-7 Trade name: XP1452

Result

No maternal effects (i.e. body weights, clinical and pathological

observations) were noted in any dose group.

3/15 rabbits miscarried in the high dose group (200 mg/kg) however, this finding was considered only bordering significance.

The number of implantations and the number and weight of the fetuses were not significantly different from the control values.

There was no difference in the distribution between male and female fetuses and there was not a significant number of malformations observed (1 fetus with aplasia of the head at 50 mg/kg and 1 fetus with internal hydrocephalus at 200 mg/kg).

No mortality was observed between the treated animals and the control

In four series of tests, the biological and biochemical parameters proved to be normal.

The histopathological results did not reveal any difference between the treated and the control animals.

Conclusion Reliability 12.12.2003

: Ultranox 626 is not consider the substance to be a teratogenic agent.

: (2) valid with restrictions

12 2003

(12)

teratogenicity in the 'Grand Fauve De Bourgogne' rabbit after oral administration, Report No. RD 3-83; in, unpublished Toxicity Summary of Ultranox 626, prepared by GE Specialty

Reimer, GJ (2003) BC Research Inc., Physical/Chemical Property of U626; CAS # 26741-

Reimer, GJ (2003) BC Research Inc., Physical/Chemical Property of U626; CAS # 26741-

Shepard, KP (1992) Toxicological Sciences Laboratory (Eastman Kodak), Bis (2,4-di-t-

butylphenyl) pentaerythritol diphosphite; Synonyms: Ultranox 626; PM 5753; skin sensitization study (Footpad method) in the Guinea Pig, HAEL No. 92-0046

53-7, Boiling Point (OECD 103), Study No. RAA6233 1257-BP

US EPA., EPIWIN v 3.10, EPI Suite Software, 2000

53-7, Hydrolytic Stability (OECD 111), Study No. RAA6233 1257-Hy

Chemicals, 1997

(13)

(14)

(15)

(16)